EAST Search History

Re f #	Hits	Search Query	DBs	Defau It Opera tor	Plur als	Time Stamp
L1	119	(548/401).CCLS.	US-PGP UB; USPAT; USOCR ; EPO; JPO; DERWE NT; IBM_T DB	OR	OFF	2007/12/09 19:05
L2	262	(548/405).CCLS.	US-PGP UB; USPAT; USOCR ; EPO; JPO; DERWE NT; IBM_T DB	OR	OFF	2007/12/09 19:06
L3	3	I1 and I2	US-PGP UB; USPAT; USOCR ; EPO; JPO; DERWE NT; IBM_T DB	OR	ON	2007/12/09 19:06

12/9/07 7:07:07 PM Page 1

EAST Search History

L4	1	l1 and oxazaborolidine	US-PGP UB; USPAT; USOCR ; EPO; JPO; DERWE NT; IBM_T DB	OR	ON	2007/12/09 19:06
L5	7	l2 and oxazaborolidine	US-PGP UB; USPAT; USOCR ; EPO; JPO; DERWE NT; IBM_T DB	OR	ON	2007/12/09 19:06

12/9/07 7:07:07 PM Page 2

10574,871> 12/09/2007

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COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 0.21 0.21

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http://www.cas.org/infopolicy.html

=> s oxazaborolidine

558 OXAZABOROLIDINE

215 OXAZABOROLIDINES

L1 605 OXAZABOROLIDINE

(OXAZABOROLIDINE OR OXAZABOROLIDINES)

=> s l1 and complexes

759694 COMPLEXES

4 COMPLEXESES

759695 COMPLEXES

(COMPLEXES OR COMPLEXESES)

L2 52 L1 AND COMPLEXES

=> s 12 and chiral

119944 CHIRAL

17 CHIRALS

119948 CHIRAL

(CHIRAL OR CHIRALS)

10574,871> 12/09/2007

L3 32 L2 AND CHIRAL

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(IN OR INS)

276067 SITU 272 SITUS 276324 SITU

(SITU OR SITUS)

131 IN(L)SITU

L40 L3 AND IN(L)SITU

=> d 13 ibib abs hitstr tot

SAEED Page 3

L1 ANSWER 1 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2007:76720 CAPLUS
DOCUMENT NUMBER: 146:337266
TITLE: Chiral Oxaraborolidine-Aluminum
Bromide Complexes Are Unusually Powerful and
Effective Catalysts for Enantioselective Diels-Alder

Effective Catalysts for Enantioselective Diels-Aider Reactions Liu, Duan: Canales, Eda: Corey, E. J. Department of Chemistry and Chemical Biology, Harvard University, Cambridge, NA, 02138, USA Journal of the American Chemical Society (2007), 129(6), 1498-1499 CODEN: JACSAT: ISSN: 0002-7863 American Chemical Society AUTHOR(S): CORPORATE SOURCE:

SOURCE:

PUBLISHER:

DOCUMENT TYPE: LANGUAGE:

Journal English CASREACT 146:337266 OTHER SOURCE(S):

AB Treatment of the chiral exaraborolidine I with AlBr3 generates the 1:1 complex II, which is an even more potent Lewis acid catalyst than protonated I for enantioselective Diels-Alder reactions. Only 4 mol % of catalyst II is required to achieve yields and enantiomeric purities of 90% over a broad range of achiral dienes and dienophiles.

ligand from which II is derived can be recovered easily and with high efficiency. The method is illustrated by 22 examples. E.g., Diels-Alder reaction of cyclopentadiene and di-Et tumerate, catelyzed by II, gave

L3 ANSWER 2 OF 32 CAPLUS COPYRIGHT 2007 ACS on STA ACCESSION NUMBER: 2007:39912 CAPLUS DOCUMENT NUMBER: 147:501894 TITLE: Asymmetric reduction of prochi

TITLE: AUTHOR(S): CORPORATE SOURCE:

MENT NUMBER: 147:501894

Asymmetric reduction of prochiral ketones

Asymmetric reduction of prochiral ketones

College of Chemistry and Materials Science, SCUFN,

Wuhan, 410074, Peop. Rep. China

CE: Zhongnan Minzu Daxue Xuebao, Ziran Kexueban (2006),

ZS(31), 27-32

CODEN: ZMOXA3; ISSN: 1672-4321

ADONGNAN HINZU DAXUE Xuebao Bianjibu

MENT TYPE: Journal; General Review

Chinese

A review. In this review, the asym. reduction of prochiral ketones and

SOURCE

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

recent advances have been described, including application of chiral oxaraborolidine catalysts, reduction by chirally modified lithium sluminum hydride reagents, application of chirally modified sodium borohydride or potassium borohydride, application of chiral phosphoramide catalysts, transfer hydrogenation of prochiral ketones catalyzed by chiral metal complexes, asym, reduction of prochiral ketones by phose-transfer catalysis, enantioselective reduction of ketones with enzymes.

L3 ANSWER 1 OF 32 CAPLUS COPYRIGHT 2007 ACS ON STN product III.
REFERENCE COUNT: 16 THERE ARE 16 CITED REFER THIS (Continued) 16 THERE ARE 16 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

L3 ANSMER 3 OF 32 CAPLUS COPYRIGHT 2007 ACS on 5TN
ACCESSION NUMBER: 2007:23509 CAPLUS
DOCUMENT NUMBER: 146:296026
TITLE: The Mg-Oppenauer oxidation as a mild method for the synthesis of aryl and metallocenyl ketones XIOetzing, Ralf J.; Krasovskiy, Arkady; Knochel, Paul CORPORATE SOURCE: Dep. Chem. Biochem., Ludwig-Maximilians-Univ.,

AUTHOR(S): CORPORATE SOURCE: Munich,

Munich,

SOURCE: Chemistry--A European Journal (2006), 13(1), 215-227
CODEN: CEUJED: ISSN: 0947-6539

PUBLISHER: Wiley-VCH Verlag GmbH & Co. KGAA

DOCUMENT TYPE: Journal
LANGUAGE: English
OTHER SOURCE(S): CASRRACT 146:296026

AB Mg alkoxides undergo a hydride-transfer oxidation with benzaldehyde as
the

oxidant. This Mg variant of the Oppenauer oxidation was used for the synthesis of polyfunctional biaryl ketones. Licl was found to promote this reaction by enhancing the solubility of Mg alkoxides. Twelve functionalized unsym. diaryl ketones were prepared from the esponding Mg alkoxides in 85% to 98% yields. E.g., 2-bromobenzophenone was obtained

of the presence of LiCl, followed by oxidation with benraldehyde.

in the presence of Lit, rollowed by condetion the selectron-accepting nor -donating substituents appear to influence the reaction course. Although one ortho-substituent does not retard the reaction, the presence of two ortho-substituents inhibited the Mg-Oppenauer reaction completely due to the high steric hindrance. This mild oxidation method was especially useful for preparing ketones

ing a metallocenyl unit as well as various new ferrocenyl ketones and tricarbonylchromium complexes. Addition of PhMgCl·LiCl to cymantrene aldehyde and oxidation with benzaldehyde afforded benzoylcymantrene in 80% yield. Various aromatic Grignard reagents were

readily added to tricarbonylchromium benzaldehyde within 10 min at -20°. Oxidation with benzaldehyde within 12 h at room temperature gave

ketones. E.g., addition of 4-Me2NC6H4MgBr·LiCl (from 4-Me2NC6H4Br, Mg and LiCl) to tricarbonylchromium benzaldehyde and oxidation with

and LiCl) to tricarbonylchromium benzaldehyde and oxidation with benzaldehyde gave 4-N,N-dimethylaminobenzoyltricarbonylchromium in 92% yield. With nonfunctionalized or electron-rich ketones, 65% to 92% yields were obtained. With halogenated arylmagnesium reagents, 20% to 59% yields

observed This last class of ketones was reduced with the CBS catalyst

Corey-Bakshi-Shibata, di-Ph oxaraborolidine) and decomplexed with iodine to give in 89% to quant. yield chiral benzhydrol complexes with high enantioselectivity enabling an asym. synthesis of electron-rich or -poor benzhydryl alcs. (up to 94% e8). E.g., reduction of

4-methoxybenzoyltricarbonylchromium benzene with borane-dimethyl sulfide complex, followed by decomplexation with iodine gave (R)-(4-methoxyphenyl)phenylmethanol in 90% yield and 93% ee.

REFERENCE COUNT: 123 THERE ARE 123 CITED REFERENCES AVAILABLE FOR

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ANSWER 3 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN (Continued)
THIS RECORD. ALL CITATIONS AVAILABLE IN THE REFORMAT
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CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
11TLE:
AUTHOR(5):
CORPORATE SOURCE:
SOURCE:
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   AB In this review, asym. reduction of processing advances have been described, including application of chiral oxazaborolidine catalysts, reduction by chirally modified lithium aluminum hydride reagents, application of chirally modified sodium borohydride or potassium borohydride, application of chiral phosphoramide catalysts, transfer hydrogenation of prochiral ketones catalysed by chiral metal complexes, asym. reduction of prochiral ketones by phase-transfer catalysia, enantioselective reduction of
                                                                                ketones with enzyme. A review.
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ANSWER 5 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN
SSION NUMBER: 2005:740583 CAPLUS
E: Enantioselective cyclopropyl ketone-based halo eldol
reactions
 ACCESSION NUMBER:
TITLE:
                          Enantisselective cyclopropyl ketone-based halo aldol reactions

IOR(S):

Li. Guigen: Timmons, Cody
Department of Chemistry and Biochemistry, Texas Tech
University, Lubbock, TX, 79409, USA

(CE: Abstracts of Papers, 230th ACS National Meeting,
Washington, DC, United States, Aug. 28-Sept. 1, 2005
(2005), ORGN-535. American Chemical Society:
Washington, D. C.
CODEN: 69MFCL

MOENT TYPE: Conference: Meeting Abstract; (computer optical disk)
BUAGE: English
Novel asym. halo aldol-type reactions have been developed using enolates
derived from cyclopropyl ketones. The enolate is formed by reacting
cyclopropyl ketones with trimethylsilyl iodide. Subsequent addition to
aldehydes is promoted by chiral oxezaborolidine
complexes, which are easily generated in situ by treating a
protected amino acid with borsen. A variety of amino acids and
lecting
groups were acreened, and N-heptafluorobutyryl-phenylalanine was found to
be optimal for controlling enantioselectivity. Calens. have been
performed to characterize the nature of the enantio-induction. In a
similar manner, sulfonyl-protected imines have given promising results
when chiral sluminum complexes are used.
AUTHOR(S):
CORPORATE SOURCE:
SOURCE
 DOCUMENT TYPE:
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L3 ANSWER 6 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2005:325360 CAPLUS DOCUMENT NUMBER: 142:392277
                                                                                                                                                                                                                                 142:392277
In situ preparation of chiral compounds derived from oxazaborolidine-borane complexes and their use as catalysts in asymmetric reductions of ketones and ether oximes Burgos, Alain; Bertrand, Blandine; Frein, Stephane; Pluvie, Jean Francois; Roussiasse, Sonia PPG-Sipsy, Fr.
Demande, 30 pp.
CODEN: FRXXBL
Patent
    TITLE:
  INVENTOR(S):
  PATENT ASSIGNEE(S):
SOURCE:
  DOCUMENT TYPE:
    LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
         PATENT NO. KIND DATE

FR 2860794 Al 20050415 FR 2003-11838 2003100.
FR 2860794 Bl 20050415 FR 2003-11838 20041011
FR 2860795 Bl 20050415 FR 2004-10701 20041011
FR 2860795 Bl 20050415 FR 2004-10701 20041011
FR 2860795 Bl 20050421 W0 2004-FR2573 20041011
W0 2005035540 A2 20050421 W0 2004-FR2573 20041011
W0 2005035540 A3 20050609
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MK, MZ, NA, NI, NO, NZ, OM, PG, PH, PI, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TH, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
RW: BW, GH, GM, KE, LS, MM, HZ, NA, SD, SL, SZ, TZ, UG, ZM, ZM, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, SI, SK, TR, BF, BJ, CF, CG, CT, CM, GA, GN, GO, GW, ML, MR, NE, SN, TD, TG

EP 1673375 Bl 20070801
R: AT, BE, CR, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, FI, RO, CY, TR, BG, CZ, EE, HU, PL, SK
CN 1867571 A 20061028 CP 2004-817157 20041011
AT 366672 T 20070405 JP 2006-5304311 20041011
AT 360672 T 30070405 JP 2006-5304311 20041011
AT 360672 T 30070405 JP 2006-5304311 20041011
AT 360672 T 30070405 JP 2006-5304311 20041011
CN 1867571
JP 2007508280
AT 368672
KR 2007026314
US 2007055068
PRIORITY APPLN. INFO.:
```

OTHER SOURCE(s): MARPAT 142:392277

AB The invention is related to the in situ preparation of chiral compds, derived from oxazaborolidine-borane complexes by reacting a metal borohydride with a Levis base, and an ester of an inorg. acid, followed by addition of an optically active maino-alc, and to their use in the preparation of chiral alcs, and ketones by asym. reduction of prochiral ketones and ether oximes. The method eliminates the use of 12 in the preparation of the oxesaborolidine-borane complex. Thus, NABH4 in THY was mixed with PhNEEZ, the mixture cooled to 5°, Me2SO4 added and the mixture stirred at 20° for 1 h, and finally mixed with

L3 ANSWER 6 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN (Centinued)

(R1-diphenylprolinol at 20° for 1 h. A soln. of

3-chloro-1-(2-thienyl)propanone in THF was added to the above preheated
maxt. over a period of 1.5 h, followed by hydrolysis for 1 h at 20°
to give the corresponding alc. in high chem. purity.

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE

L3 ANSWER 7 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2004:1053291 CAPLUS
DOCUMENT NUMBER: 143:30591;
Synthesis of chiral amino alcohols by enantioselective borane reduction
AUTHOR(S): Shen, Zong-xuan; Li, Yong-hua; Qin, Hong-bing; Zhang, Ya-wen

CORPORATE SOURCE:

Ya-wen Key Lab of Organic Synthesis of Jiangsu Province, College of Chemistry and Chemical Engineering, Suzhou University, Suzhou, Jiangsu, 215006, Peop. Rep. China Huaxue Yanjiu (2004), 15(3), 23-26, 49 CODEN: HUAYF4; ISSN: 1008-1011 Huaxue Yanjiu Bianjibu

PUBLISHER:

DOCUMENT TYPE; Journal LANGUAGE:

OTHER SOURCE(S):

MENT TYPE: Journal MARKET TYPE: Chinese Chinese R SOURCE(S): Chinese R SOURCE(S): CASREACT 143:305917

A group of α - and β -dialkylamino ketones were reduced enantiosalectively by using BB3-THF and the in-situ formed chiral oxazaborolidine. The formed optically active aminoalc.-borane complexes, which, after deboration gave optically active amino alcs. with high enantiomeric excess. The effects of the substrate structure on the enantioselectivity of the reaction are also discussed.

L3 ANSWER 8 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2004:531764 CAPLUS DOCUMENT NUMBER: 141:243031 Highly Stereoselective Approach Alk-2-yne-1,4-diols

Highly Stereoselective Approach to by Oxazaborolidine-Mediated Reduction of

AUTHOR (S):

by Oxazaborolidine-Hediated Meduction of Alk-2-yne-1,4-diones Ariza, Xavier: Bach, Jordi: Berenguer, Ramon; Farras, Jaume: Fontes, Hontserrat: Garcia, Jordi: Lopez, Marta: Ortiz, Jordi Departament de Quimica Organica, Universitat de Barcelona, Barcelona, E-08028, Spain Journal of Organic Chemistry (2004), 69(16),

CODEN: JOCEAH: ISSN: 0022-3263 American Chemical Society Journal English CASREACT 141:243031

PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
OTHER SOURCE(S):

CORPORATE SOURCE:

SOURCE: 5307-5313

The authors performed borane-mediated reduction of a series of sym. alk-2-yne-1.4-diones, e.g. I, in the presence of a chiral oxezaborolidine to afford (R.R.)-alk-2-yne-1.4-diols, e.g. II, in good yields and high stereoselectivities (up to 99.9% ee). In some

the stereochem. purity of II was improved by a two-step process: (i) temporary transformation of II into its vic-dibromo derivs. which allowed the removal of the minor mean isomer by chromatog., and (ii) regeneration of the enanticentiched diols II with SmIZ. Reduction of the hexacarbonyldicebalt complexes derived from sym. alk-Z-yne-1,4-diones, e.g. I, was also successful.

REFERENCE COUNT: 57 THERE ARE 57 CITED REFERENCES AVAILABLE FOR THIS

FORMAT

RECORD. ALL CITATIONS AVAILABLE IN THE RE

L3 ANSWER 9 OF 32 CAPLUS COPYRIGHT 2007 ACS ON STN
ACCESSION NUMBER: 2003:560200 CAPLUS
DOCUMENT NUMBER: 139:260801
TITLE: Quantum Hechanical Study of St 139:260801 Quantum Mechanical Study of Stereoselectivity in the Oxazaborolidine-Catalyzed Reduction of

AUTHOR(S):

CORPORATE SOURCE:

SOURCE:

PUBLISHER:

DOCUMENT TYPE: LANGUAGE:

LE: Quantum Mechanical Study of Stereoselectivity in the Oxazaborolidine-Catalyzed Reduction of Acetophenone
HOR(S): Alagona, Gulliano: Ghio, Caterina: Persico, Maurizio; Tomasi, Simone
PORATE SOURCE: Istituto per i Processi Chimico-Fisici. CNR, Pisa, I-56124, Italy
GOMEN: Journal of the American Chemical Society (2003), 125(13), 1027-10039
CODEN: JACSAT: ISSN: 0002-7863
LISHER: American Chemical Society
UMENT TYPE: Journal
GUAGE: English
Chiral oxazaborolidines, known as CBS catalysts after
the work of Corey, Bakshi and Shibata, are used for the stereoselective reduction of prochiral ketones to secondary chiral alcs. Due to their relative low cost, ease of use, and high selectivity, their popularity has remarkably grown in the last 15 yr. Oxazaborolidine
-catalyzed redns. have been much studied, both exptl. and putationally,
by means of semiempirical methods. Though, a more accurate high level quantum mech. study on the complete system, capable of elucidating reliably the origins of stereoselectivity, is still lacking. Therefore, the acetophenone (PhNK) reduction with Corey's oxazaborolidine has been modeled for the first time with ab initio and DFT-B3LFY calcns. on the complete system as well as with AMI. Calcns. on the complexation of BH3 to CBS, which can occur only in a cis fashion with respect to the hydrogen on the Stereogenic C-4 carbon atom, have allowed us to confirm the great rigidity of Corey's catalyst, possibly determining its ellent
enanticselectivity. Acetophenone-CBS-BH3 complexes were characterized at various levels of theory, and it was found that the picture obtained depends heavily on the method adopted. A computational strategy for identifying the hydride transfer transition states of the competing pathways was developed and teated, using a model system for which the transition state geometry was already known. The application, the TS search method to the reduction of acetophenone allowed the

the TS search method to the reduction of acetophenone allowed the characterization of the TS's for the competing pathways in this reaction making it possible to predict with good quant. accuracy the stereochem. outcome of the reaction at all the levels of theory adopted. The characterization of the intermediate oxazadiboretane products confirmed that the highly exothermic hydride transfer provides the thermodynamical drive for the reaction.

REFERENCE COUNT: 78 THERE ARE 78 CITED REFERENCES AVAILABLE FOR THIS

FORMA1

RECORD. ALL CITATIONS AVAILABLE IN THE RE

L3 ANSWER 10 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2002:906244 CAPLUS
DOCUMENT NUMBER: 138:4202
TITLE: Preparation of iron-group metal-bound
oxaraborolidines as reusable catalysts for
enantioselective reduction of ketones
INVENTOR(5): Court. Jean! Long. Moniour.

INVENTOR (S): PATENT ASSIGNEE (S): Court, Jean: Lopez, Monique Centre National de la Recherche Scientifique CNRS,

Fr. SOURCE: PCT Int. Appl., 19 pp. CODEN: PIXXD2

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

P.	ATENT															ATE	
W	2002	0948	37		A1		2002	1128		WO	2002	- FR1	602		2	0020	513
	W:	ΑE,	AG,	AL.	AM,	AT,	AU,	AZ,	BA,	ВВ	, BG	, BR	, BY,	BZ,	CA,	CH,	CN,
		co,	CR,	Cυ,	CZ,	DE,	DK,	DM,	DZ.	EC	, EE	, Es	, FI,	GB.	GD,	GE,	GH,
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	RW:											. UG	, zm,	2W.	AT.	BE.	CH.
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	5 2004									υs	2003	-476	668		2	0031	104
U	S 6825	370			B2		2004	1130									
PRIORI	TY APP	LN.	INFO	. :						FR	2001	-661	2		A 2	0010	518

OTHER SOURCE(5): CASREACT 138:4202; MARPAT 138:4202

AB The invention concerns oxazaborolidine compds. fixed on Raney

Ni, Raney Co or Raney Fe, the method for preparing same, and the use of compds. as reusable catalysts for enantioselective reduction of ketones

produce chiral alcs. For example, Raney Ni was treated in a suspension of THF with LiBH4 followed by BH3·(N,N-diethylaniline) or another borane adduct to give NiSB. An amino alc. (e.g. (15,ZR)-(+)-2-amino-1,2-diphenylethanol) in THF was added to the NiSB suspension to give the catalyst. Borane adduct is added to the catalyst and then the ketone in THF is reduced in its presence. P-fluoroacetophenone was reduced with 93% ee when the catalyst was used for the lst time, followed by 91% and 90%, resp. for the 2nd and 3rd

L3 ANSWER 11 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:902346 CAPLUS DOCUMENT NUMBER: 138:15364 Asymmetric Synthesis of anti-a

SAEED

138:153636
Asymmetric Synthesis of anti- and syn-A-Amino
Alcohols by Reductive Cross-Coupling of Transition
Metal-Coordinated Planar Chiral
Arylaldehydes with Aldimines
Tanaka, Yoshie; Taniguchi, Nobukazu, Kimura,

WO 2002-FR1602

w 20020513

AUTHOR(S): Takayuki;

CORPORATE SOURCE:

Uemura, Motokazu
Department of Chemistry, Faculty of Integrated Arts
and Sciences, Osaka Prefecture University, Osaka,
599-8511, Japan
Journal of Organic Chemistry (2002), 67(26),

SOURCE: 9227-9237

CODEN: JOCEAH: ISSN: 0022-3263 American Chemical Society Journal English CASREACT 138:153636 PUBLISHER:

DOCUMENT TYPE: LANGUAGE:

OTHER SOURCE(S):

Sml2-mediated cross-coupling of N-(tosyl)ferrocenylideneamine with planar chiral ferrocenecarboxaldehydes or benzaldehyde Cr tricarbonyl complexes gave diastereoselectively the corresponding anti-fi-anino alc. derivs. in good yields, while N-(tosyl)benzylideneamine produced syn-fi-amino alcs. by coupling with planar chiral arylaldehydes. E.g., Sml2 mediated reaction of (*)-(S)-c-lodoferrocenecarboxaldehyde with FcCH:NTs in THF (Fc = ferrocenyl) gave anti-fi-amino alc. derivative I (R = p-MacGH8502) in 94% yield with 94% ee while Sml2-mediated cross-coupling of PhCH:NTS with Cr(CGH861CO3) gave syn-fi-amino alc. PhCHNMTs(CHOAP)Ph with high diastereoselectivity (syn/anti = 97/3) in 73% overall yield after acctylation and subsequent demetalation. Dynasic kinetic resolution of a configurationally equilibrated reactive species generated from achiral N-(tosyl) ferrocenylideneamine and -benzylideneamine by reduction with SmI2

Sm12 was observed in the cross-coupling with planar chiral arylaidshydes giving both antipodes of B-amino alcs. depending on the planar chirality. The obtained anti-B-amino alc, with the ferrocene ring was used as a chiral ligand for catalytic asym. reduction of acetophenone. E.g., catalytic asym. reduction of acetophenone in presence the

oxazaborolidine formed in situ from anti-β-amino alc. II, MeB(OH)2, and B2H6 gave (S)-PhCH(OH)Me in 83% yield and 91% ee. A

L3 ANSWER 10 OF 32 CAPLUS COPYRIGHT 2007 ACS on STM (Continued)
times. Product sepn. is easier using this system than when the catalysts
are bound to org. polymers.

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE

ANSWER 11 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) reaction mechanism was proposed to rationalize the obsd.

stereoselectivity of the cross-coupling of N-(tosyl)ferrocenylideneamines with arylaldehydes. The efficient achievement of cross-coupling between ferrocenecarboxaldehyde and N-(arylaulfonyl)ferrocenylidenimines was attributed to different redn. potentials between both substrates.

REFERENCE COUNT: 94 THERE ARE 94 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

PUBLISHER:

L3 ANSWER 12 OF 32 CAPLUS COPYRIGHT 2007 ACS On STN
ACCESSION NUMBER: 2002:616524 CAPLUS
TITLE: Predictive modeling of chemical reactions
Soltzberg, L. J.; Lee, Nancy E.: Honda, Ayako;
Sanford, Micheller Sekoni, Mojisola
CORPORATE SOURCE: Department of Chemistry, Simmons College, Boston, MA, 02115, USA

SOURCE:

02115, USA Abstracts of Papers, 224th ACS National Meeting, Boston, MA, United States, August 18-22, 2002 (2002), COMP-288. American Chemical Society: Washington, D.

CODEN: 69CZPZ

DOCUMENT TYPE:

COURT: OPERFY
MENT TYPE: Conference: Heeting Abstract
UAGE: English
We have used off the shelf software (Spartan Pro) to model the outcomes

a variety of chemical reactions. Geometries of the reactant mols. are

optimized, and the mols, are given an initial orientation that might correspond to a reactive collision. Geometry optimization is then run on the assembly. Application of this procedure with a low level ab initio Hamiltonian has replicated the correct products and relative reactivities for the reduction of hydroquinone by various alkylammonia borane complexes. With a semi-empirical Hamiltonian, application of this method to the oxazaborolidine-catalyzed reduction of diverse ketones gives not only the correct products but reproduces the enantioselectivity of these redns, when carried out with a chiral catalyst.

L3 ANSWER 13 OF 32 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 2002:585820 CAPLUS DOCUMENT NUMBER: 137:278751
TITLE: Broad-Space....

137:278751
Broad-Spectrum Enantioselective Diels-Alder Catalysis by Chiral, Cationic Oxaraborolidines
Ryu, Do Hyun: Lee, Thomas W.; Corey, E. J.
Department of Chemistry and Chemical Biology, Harvard
University, Cambridge, NA, 02138, USA
Journal of the American Chemical Society (2002),
124(14), 9992-9993
CODEN: JACSAT; ISSN: 0002-7863
American Chemical Society
Journal AUTHOR(S): CORPORATE SOURCE: SOURCE:

DOCUMENT TYPE: LANGUAGE: Journal

English CASREACT 137:278751 OTHER SOURCE(S):

The cationic chiral Lewis acids I (R = Ph, 3,5-Me2C6H3), generated by triflic acid protonation of the corresponding neutral oxazaborolidines, serve as excellent catalysts for Diela-Alder addition of cyclopentadiene to a wide variety of dienophiles. Adducts

have been obtained in excellent yield and enantioselectivity from u.f.-unastd. esters. lactones, and cyclic ketones. The absolute facial selectivity for each of these substrates follows a common patter which differs from that observed with u.f.-enals. The different reaction channels can be understood in terms of pathways via complexes (for u.f.-enals) and (for u.f.-ences and esters).

REFERENCE COUNT: 23 THESE ARE 23 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L3 ANSWER 14 OF 32
ACCESSION NUMBER:
DOCUMENT NUMBER:
136:385883
Asymmetric Diels-Alder Reactions Catalyzed by a Triflic Acid Activated Chiral Oxazaborolidane
AUTHOR(S):
CORPORATE SOURCE:

SOURCE:

SOURCE:

DOCUMENT TYPE:
LANGUAGE:
DOCUMENT TYPE:
LANGUAGE:
DOCUMENT TYPE:
LANGUAGE:
OTHER SOURCE(S):
CASREACT 136:38583
CASREACT 136:38583

Oxazaborolidines I (R = 2-Mec6H4, 3,5-Me2C6H3; R1 = 2-Mec6H4), prepared from proline-derived a.a-diarylpyrrolidinemethanols and boroxines upon reflux in benzene, are activated by triflic acid to give powerful Lewis acid catalysts; in the presence of 6 mol's of the triflic acid-activated oxazaborolidines at -95°, the asym. blela-Alder cycloaddns. of cyclic and acyclic dienes with either 2-Me or 2-bromoacrolein occur to give monocyclic or bicyclic cyclohexencerboxaldehydes in 58-981 yields and in 91-971 ee. E.g., (S)-a,a-bis(3,5-dimethylphenyl)pyrrolidine-2-methanol was attirred in toluene with 1,3,5-tri(2-tolyl)boroxine and heated at 60° for 4 h followed by distillation to give I (R = 3,5-Me2C6H3; R1 = 2-MeC6H4) as a clear oil. E.g., 1.0 equivalent of triflic acid was a colution of I (R = 3,5-Me2C6H3; R1 = 2-MeC6H4) as a clear oil.

I to a solution of I (R = 3,5-Me2C6H3; Rl = 2-MeC6H4) in methylene chloride; the solution of I (R = 3,5-Me2C6H3; Rl = 2-MeC6H4) in methylene chloride; the solution was cooled to -95° and 2-bromoacrolein and 1,3-butadiene were added dropwise through a cannula; after stirring for 2 h at -95°, the mixture was warned to room temperature and worked up to give cyclohexenecarboxaldehyde II in 98% yield and 97% ee. NMR studies of I

Ph; Rl = Me) in the presence of triflic acid showed that an
equilibrium mixture
 of triflic acid-oxazaborolidine complexes exists; the
 species equilibrate slowly under the reaction conditions and more rapidly
 at 0°. The catalysts generated by treatment of I with triflic acid
 are estimated to be of similar acidity to triflic acid catalysts
generated
 from oxazaborolidines and methanesulfonic acid were ineffective
 at promoting Diels-Alder cycloaddn. reactions, allowing one to estimate
the

acid strength of the triflic acid complex of I. I were optimized for substitution at boron and at the α -position of the

L3 ANSWER 14 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) pyrrolidinemethanol moieties. The reactions are simple to conduct, reproducible, and economical, since only ca. 6 mol % of I is required; in addn., the chiral pyrrolidinemethanol precursors of I are readily recovered for reuse (>95% efficiency) and com. available. The reaction is effective with both cyclic and acyclic dienes. A mechanism accounting for the abs. and relative stereochem. of the Diels-Alder products is proposed.

REFERENCE COUNT: 19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR

19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR

FORMAT

RECORD. ALL CITATIONS AVAILABLE IN THE RE

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L3 ANSWER 15 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2001:877643 CAPLUS
DOCUMENT NUMBER: 136:278901
ASYMMETRIC reduction of amino ketones with borane and chiral oxazaborolidine catalyst
Chiral oxazaborolidine catalyst
Zhang, Ya-Men; Shen, Zong-Xuan; Cin, Hong-Bing; Li, Yong-Hua: Yu, Kai-Bei
Department of Chemistry and Chemical Engineering, Suxhou University, Suzhou, 215006, Peop. Rep. China Chinese Journal of Chemistry (2001), 19(11),
```

CODEN: CJOCEV; ISSN: 1001-604X Science Press Journal English CASREACT 136:278901

PUBLISHER: DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S):

England

R SOURCE(S): CASREACT 136:278901

Some a-(dialkylamino) ketones and ß-(dialkylamino) ketones were reduced stereoselectively by 2 mol of borane-tetrahydrofuran in the presence of 10 molt of an in situ-formed chiral coxraborolidine, followed by diluted hydrochloric acid. The catalysts in this study included (1aR)-tetrahydro-1-mathyl-3, 3-diphenyl-1H, 3H-pyrrolo[1,2-c][1,3,2]oxaraborole, (3aS)-tetrahydro-1-methyl-3,3-diphenyl-1,3,2-oxaraborolidine. The resulting amino alc.-borane complexes were treated with hydrogen chloride-glycol-methanol to give the optically active amino alc. with the ee up to 99%. For example, the reduction of 3-(4-morpholinpyl)-1-propanone gave (+)-(aR)-a-phenyl-4-morpholinepropanol in 87.3% yield. The intermediate a-phenyl-4-morpholinepropanol-borane complex was characterized by x-ray crystallog.

RENCE COUNT: 14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR

REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L3 ANSWER 16 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2000:536261 CAPLUS
DOCUMENT NUMBER: 14:147384

An efficient synthesis of (15, 2R)-1-amino-2-indanol,
a key intermediate of HIV protease inhibitor,
indinavir

AUTHOR(S): Demir, Ayhan S., Aksoy-Cam, Hilal: Camkerten,
Nurettin: Hamamci, Haluk; Doganal, Fatos
Department of Chemistry, Middle East Technical
University, Ankara, 06531, Turk.

SOURCE: Turkish Journal of Chemistry (2000), 24(2), 141-146
CODEN: TJCHE3; ISSN: 1300-0527

PUBLISHER: Scientific and Technical Research Council of Turkey
DOCUMENT TYPE: Journal
LANGUAGE: English
CTHER SOURCE(S): CASREACT 134:147384

A (15, 2R)-1-amino-2-indanol, a key component of HIV protease inhibitor was
prepared in four steps starting from indanone. The Mn(OAcl)3 mediated
acetoxylation of indanone followed by fungus catalyzed hydrolysis of
acetoxylandanone furnished optically pure a-hydroxylandanone.
Formation and enanticselective reduction of oxime ether of
2-hydroxylandanone
afforded (15, 2R)-1-amino-2-indanol in 978 cis selectivity. 2-hydroxyindanone afforded (15,2R)-1-amino-2-indanol in 97% cis selectivity. REFERENCE COUNT: 58 THERE ARE 58 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 17 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2000:188699 CAPLUS
DOCUMENT NUMBER: 133:89130
TITLE: Nondynamic Kinetic Resolution of Configurationally

Nondynamic Kinetic Resolution or Configuration Stable Biaryl Lactones by Reduction with Oxazaborolidine-Activated Borane: AMI Studies and Experimental Verification Bringmann, Gerhard: Hinricha, Juergen; Kraus,

AUTHOR (S) :

CORPORATE SOURCE:

Wuzik, Andreas; Schulz, Tanja Institut fuer Organische Chemie, Universitaet Wuerzburg, Wuerzburg, D-97074, Germany Journal of Organic Chemistry (2000), 65(8), 2517-2527 CODEN: JOCEAN; ISSN: 0022-1263 American Chemical Society Journal English CASREACT 133:89130 SOURCE:

PUBLISHER:

DOCUMENT TYPE: LANGUAGE:

OTHER SOURCE(\$):

t-Bu 11

The complete mechanistic course of the atroposelective ring opening of a lactone-bridged biaryl, dinaphth(2,1-c:1',2'-e]oxepin-1-(5H)-one (3 = I), with a chiral oxazaborolidine-BH3 complex was calculated using the semiempirical AMI method. The first hydride transfer to the activated carbonyl function of the adduct complexes was alsorated to be the selectivity-determining step in the postulated step

mechanism. The calculated enantioselectivity is in good accordance with

exptl. results, so that related calons, were performed on the atroposelective ring opening of a sterically strongly hindered and therefore also configurationally stable six-membered biaryl lactone, 1,3-di-tert-butyl-64*-benzo[b]naphtho[1,2-d]pyran-6-one (6f = II). These calons, predicted a highly (M)-selective reduction of 6f (kM/kP = 358 at

*C), which, after the smooth preparation of 6f by intramol. biaryl coupling in high yields, was fully confirmed exptl. (hM/kP > 200 at -78 *C). Isolation of the intermediate hydroxy aldehyde (H)-14 ([M)-111, R = CMO] at the beginning of the reaction with the same enantiomeric excess as found for the corresponding alc. (M)-7f [(M)-III,

R

- CH2OH) conclusively showed the first hydride transfer step to
determine the
selectivity of this process. The good agreement of computationally
predicted and exptl. confirmed values proves the suitability of the AMI
method for mechanistic studies on even such complex reactions and opens

ANSWER 17 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) most efficient overall synthesis of sterically highly hindered biaryls,

excellent chem. (for the ring closure) and optical (for the ring

yields and for any desired exial configuration.
REFERENCE COUNT: 53 THERE ARE 53 CITED REFERENCES AVAILABLE FOR

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L] ANSMER 18 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:
DOCUMENT NUMBER:
11:31:310376
Catalytic enantioselective reactions. Part 16.
OXATABOROLIGINE extet acetals
AUTHOR(S):
CORPORATE SOURCE:
COUNCE:
COUNCE:
DOUBLISHER:
PUBLISHER:
PUBLISHER:
PUBLISHER:
ROYAL SOURCE:
CAPPORATE SOURCE:
PUBLISHER:
ROYAL SOURCE:
ROYAL SOURCE:
PUBLISHER:
ROYAL SOURCE SOURCE:
ROYAL SOURCE SOURCE:
PUBLISHER:
ROYAL SOURCE SOURCE

DOCUMENT TYPE: LANGUAGE:

English CASREACT 131:310376 OTHER SOURCE(S):

AB Asym. redns. of a-keto acetals, e.g., PhCOCH(OEt)2, using various oxazaborolidines, e.g., I (R = H, Me, Bu, Ph), and borsne reagents, e.g., BH3-THF, PhNEt2-BH3, as catalysts and hydride sources, resp., were compared. The reduction catalyzed by reagents | (R = H, Me) with N-phenylamine-borane reagents provided chiral analysis and catalysed section (R = H, Me) with N-phenylamine-borane reagents provided chiral analysis are catalysis of the section of

RECORD. ALL CITATIONS AVAILABLE IN THE RE

L3 ANSWER 20 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1996:705596 CAPLUS DOCUMENT NUMBER: 126:8173

AUTHOR/S):

126:8173
Righly Enanticenriched Propargylic Alcohols by Oxazaborolidine-Mediated Reduction of Acetylenic Ketones
Bach, Jordi: Berenquer, Ramon: Garcia, Jordi: Loscertales, Teresa: Vilarrasa, Jaume
Departament de Quimica Organica, Universitat de Barcelona, Barcelona, 08028, Spain
Journal of Organic Chemistry (1996), 61(25), CORPORATE SOURCE:

SOURCE: 9021-9025

CODEN: JOCEAH; ISSN: 0022-3263 American Chemical Society Journal English CASREACT 126:8173

PUBLISHER: DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): GI

AB An efficient and general approach to highly enantiomerically enriched propargylic ales., e.g., I (R = PhCH2CH2), by borane-mediated, oxazaborolidine, e.g., II, catalyzed reduction of ketones, e.g., PhCH2CH2COCH.tplbond.CSiMeJ, is described.

REFERENCE COUNT: 63 THERE ARE 63 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 19 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1997:338202 CAPLUS
DOCUMENT NUMBER: 127:50692
TITLE: Chiral increases

127:50692
Chiral intramolecular amine-borane
complexes as reducing agents for prochiral
ketones
Toumelin, Jean-Brice Le: Baboulene, Michel
Laboratoire des IMRCP, UMR 5623 (CNRS), Universite P.
Sabatier, Toulouse, 31062, Pr.
Tetrahedron: Asymmetry (1997), 8(8), 1259-1265
CODEN: TASYE3; ISSN: 0957-4166
Elsevier
Journal
English AUTHOR(S): CORPORATE SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S):

English CASREACT 127:50692

AB A new family of chiral amine-borane complexes, the
N-spiroazaborolidines I, were synthesized from reaction of
allylaminoethanol II (e.g., Rl = H, Me, R2 = Me, Ph) with R3B(OH)2 (R3 =
Me, Ph, H) followed by cyclization with borane dimethylsulfide. I are
stable, convenient to use, and are excellent reducing agents of prochiral
ketones (yieldreduction ≥ 95h). However, poor enantioselectivity
was obtained (ee ≤ 38h). The configuration of these mols. (cis
position between Baza and the substituent on the Boxaza), unfavorable for
a good approach of the ketone, is a possible explanation. These results
show the importance of the stereochem. of the N atom in amine-borane
complexes for asym. synthesis.

REFERENCE COUNT: 30 THERE ARE 30 CITED REFERENCES AVAILABLE FOR
THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

II

FORMAT

L3 ANSWER 21 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1994:655858 CAPLUS
DOCUMENT NUMBER: 121:255858
A combined synthetic and ab initio study of chiral oxazaborolidines structure and enantioselectivity relationships
AUTHOR(S): Quallich, George J.; Blake, James F.; Woodall, Teresa

CORPORATE SOURCE: Central Research Division, Pfizer Inc., Groton, CT,

O6140, USA
Journal of the American Chemical Society (1994),
116(19), 8516-25
CODEN: JACSAT; ISSN: 0002-7863 SOURCE:

Journal DOCUMENT TYPE:

DOCUMENT TYPE: JOURNAL LANGUAGE: English
OTHER SOURCE(s): CASREACT 121:255850
AB Investigations into the relationship of oxazaborolidine
structure to the enantioselectivity obtained in the reduction of

niral ketones revealed the intrinsic power of the mol. recognition element in the catalytic reduction This mol. recognition, two-point binding of

the catalytic reduction This mol. recognition, despending the borane
and the ketonic oxygen atom by the oxazaborolidine, assembles a trimol. complex which provides high enantiomeric excess. Enantiomeric excess was demonstrated to be dependent on the extent to which one oxazaborolidine face was precluded from attaining two-point binding and on nonbonded interactions that developed during formation of the borane-oxazaborolidine complex. As a result, erythro-aubstituted oxazaborolidines were demonstrated to be useful catalysts for enantioselective reduction of prochiral ketones. Ab initio MO calons, have been used to locate possible complexes and transition state assemblies that correspond to catalyst-borane and the

trimol. complex on a proposed reduction pathway. Geometry optimizations

carried out at the 3-21G, 6-31G(d), and MP2/6-31G(d) levels of theory. Correlation energies were computed via Moeller-Plesset perturbation theory

to the second order (MP2). Relative activation energies establish correctly the observed enantioselectivity of the two best oxszaborolidine catalysts in this study. Addnl, the diminished enantioselectivity of N-methyl-substituted catalysts was traced to conformational changes in the exo transition state. Though the relative energies obtained from the various levels of theory are similar, absolute complexation and activation energies are found to vary considerably with the level of theory employed. The existence of key intermediates was found to depend on the level of theory.

10574,871>

L3 ANSWER 22 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1994:508893 CAPLUS DOCUMENT NUMBER: 121:108893

121:108893
Quantum chemical modeling of chiral catalysis. Part 18. Conformational studies on chiral N-sulfonylated 1,3,2-oxazaborolidines and related aldehyde complexes potentially involved in the catalytic asymmetric Diels-Alder reactions Nevelainen. Vesa Div. Org. Chem., Univ. Helsinki, Helsinki, SF-00014, Finland

AUTHOR(S): CORPORATE SOURCE: riniand Tetrahedron: Asymmetry (1994), 5(4), 767-72 CODEN: TASYE3: ISSN: 0957-4166 SOURCE:

DOCUMENT TYPE: Journal LANGUAGE:

AB Properties of Lewis acidic oxazaborolidines were investigated by means of ab initio MO methods [RHF, N-sulfonylated 1,3,2-oxazaborolidine [I] as a model]. Energies of the coordination of aldehydes to oxazaborolidines were estimated by using formaldehyde and acrolein adducts of I as models. Energies (NPZ/6-31G/6-31G) of the coordination of formaldehyde to I were determined The corresponding energies of the formation of acrolein adducts, were higher.

12/09/2007

L3 ANSWER 23 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1994:492602 CAPLUS 121:22502 Convenient procedures for the asymmetric reductions utilizing α, α -diphenylpyrrolidinemethanol and borane complexes generated using the 12/NaBH4 system AUTHOR(S): Periasamy, Mariappan; Kanth, J. V. Bhaskar; Prasad,

AUTHOR(S):

 Bhenu
 Chem., Univ. Hyderabad, Hyderabad, 500 134, CORPORATE SOURCE:

India SOURCE:

Tetrahedron (1994), 50(21), 6411-16 CODEN: TETRAB: ISSN: 0040-4020 Journal English CASREACT 121:82602

DOCUMENT TYPE:

LANGUAGE: OTHER SOURCE(S): GI

Prepns. of oxazaborolidine in situ in benzene using u.a-di-phenylpyrrolidinemethanol and diborane, generated from the I-Na borohydride system are described. The oxazaborolidine (10 mol 1), generated by the reaction of u.a-diphenylpyrrolidinemethanol and diborane in benzene followed by heating with N.N-diethylaniline, in combination with borane-THF complex reduces acctophenone to 1-phenylethanol in 94.71 ee. Formation of the oxazaborolidine complex I was facilitated in the presence of an amine.

L3 ANSWER 24 OF 32 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 1994:409461 CAPLUS DOCUMENT NUMBER: 121:9461

TITLE:

AUTHOR(S): CORPORATE SOURCE:

SOURCE: DOCUMENT TYPE:

SSION NUMBER: 1994:409461 CAPLUS
MENT NUMBER: 121:9461
E: Quantum-chemical modeling of chiral
catalysis. Part IS. On the role of hydride-bridged
borane-alkoxyborane complexes in the
catalytic enantioselective reduction of ketones
promoted by chiral oxazaborolidines

ORATE SOURCE: Div. Org. Chem., Univ. Helsinki, Helsinki, SF-00014,
Finland
CE: Tetchedron: Asymmetry (1994), 5(2), 289-96
CODEN: TASYES; ISSN: 0957-4166
MENT TYPE: Journal
MUMGE: English
Hydride-bridged borane-alkoxyborane complexes of
oxazaborolidines were investigated by means of ab initio MO
methods (RHF). The complexes were found to be stable in the
absence of Lewis basic solvents. In the presence of a Lewis basic
ent

ent
the borane-alkoxyborane complexes were predicted to decompose
leading to the formation of borane-solvent complexes of the
oxazaborolidines. A new class of oxazaborolidine
catalysts, of which the mechanism of regeneration would be based on
borane-borane hydride exchange, was invented.

L3 ANSWER 25 OF 32 CAPLUS COPYRIGHT 2007 ACS ON STN
ACCESSION NUMBER: 1994:217770 CAPLUS
DOCUMENT NUMBER: 120:217770
TITLE: Quantum Charles

120:217770

Quantum chemical modeling of chiral
catalysis. Part 13. Role of borane O-adducts in the
enantioselective reduction of ketones catalyzed by
chiral oxazaborolidines
Nevalainen, Vesa
Div. Org. Chem., Univ. Helsinki, Helsinki, 00014,
Finland
Tetrahedron.

AUTHOR(S): CORPORATE SOURCE:

SOURCE:

Tetrahedron: Asymmetry (1993), 4(9), 2001-10 CODEN: TASYE3; ISSN: 0957-4166

DOCUMENT TYPE: LANGUAGE: English

Plausible reactions of Lewis bases (ketones and ethers) with borane O-adducts of chiral oxazaborolidines, e.g., I, used as catalysts in the enantioselective reduction of ketones were investigated

means of ab initio MO methods. Properties of the O-adducts were found to be different from those of the corresponding N-adducts. The O-adducts were not able to form complexes with ketones and ethers similar to those of the corresponding N-adducts proposed to be essential for the performance of oxazaborolidines as chiral catalysts in the enantioselective reduction of ketones.

10574,871>

L3 ANSWER 26 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
1993:670371 CAPLUS
119:270371 Quantum chemical modeling of chiral
catalysis. Part 12. Influence of the nature of the
ring system on binding in ketone-borane
complexes of chiral
oxazaborolidines used as catalysts in the
enantioselective reduction of ketones
AUTHOR(S):
Newslainen, Vesa
Div. Org. Chem., Univ. Helsinki, SF-00014, Finland
Tetrahedron: Asymmetry (1993). 4(7), 1597-602
CODEN: TASVEJ; ISSN: 0957-4166
DOCUMENT TYPE:
DOCUMENT TYPE:
LANGUAGE:
DOCUMENT TYPE:
DOCUMENT TYPE:
DOCUMENT TYPE:
DOCUMENT TYPE:
Oxerable of the stones (formaldehyde as model) to borane N-adducts of
1-aza-2-bora-3-oxbicyclol3.3.0)octane (1) and 1,3,2oxazaborolidine was investigated by means of ab initio MO methods.
Coordination energies were more pos. in the case of the more strained and
rigid bicyclic I. Why I derivs. can perform better as catalysts than the

12/09/2007

ACCESSION NUMBER:
DOCUMENT NUMBER:
1993:670321 CAPLUS
1992:70321 C

as a model of carbonyl compds, and an adduct of formaldehyde to N-sulfonyl-1,3,2-oxaraborolidine as a model of the complexes. In this six-ring system, the oxygen of formaldehyde was bound to the boron of oxaraborolidine and the carbon to one of the oxygens of the N-sulfonyl group. As a consequence of this unusual complexation, the C:O double bond of formaldehyde lengthened substantially; the plenar shape of the carbonyl was distorted and the

charge of the carbonyl carbon increased about 100%.

L3 ANSWER 28 OF 32 CAPLUS COPYRIGHT 2007 ACS ON STN ACCESSION NUMBER: 1993:234104 CAPLUS DOCUMENT NUMBER: 118:234104

TITLE:

118:234104
Quantum chemical modeling of chiral
catalysis. Part 8. The conformational freedom of the
ketone of ketone-borane complexes of
oxazaborolidines used as catalysts in the
enantioselective reduction of ketones
Newalainen, Vesa
Dep. Chem. Univ. Helsinki, Helsinki, SF-00100,
Finland
Tetrahadron.

AUTHOR(S): CORPORATE SOURCE:

SOURCE:

DOCUMENT TYPE:

LANGUAGE:

Finland
Tetrahedron: Asymmetry (1992), 3(12), 1563-72
CODEN: TASYES: ISSN: 0957-4166

MENT TYPE: Journal
SUAGE: English
Standard ab initio MO methods were employed to study conformational

dom of
the ketone of ketone-borane complexes of chiral
oxazaborolidines used as catalysts for the enantioselective reduction
of ketones. A formaldehyde-borane complex of 1,3,2oxazaborolidine was used as a model system. A new conformation
was found which was energetically more advantageous than the original one
predicted by Corey et al. The new conformation was predicted to be
destabilized by bulky substituents at the C-5 of the ring. A new class

potential oxazaborolidine catalysts for the enantioselective reduction of ketones was found.

L3 ANSWER 29 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1992:425621 CAPLUS
DOCUMENT NUMBER: 117:25621
TITLE: Quantum chemical modeling of chiral
catalysis. Part 4. On the hydride transfer in

ketone

complexes of borane adducts of oxazaborolidines and regeneration of the

oxataborolidines and regeneration of the catalyst Nevalainen, Vesa Dep. Chem., Univ. Helsinki, Helsinki, SF-00100, Finland Tetrahedron: Asymmetry (1991), 2(11), 1133-55 CODEN: TASYE3; ISSN: 0957-4166 Dournal English AUTHOR(S): CORPORATE SOURCE:

SOURCE:

DOCUMENT TYPE:

LANGUAGE:

Hydride transfer and regeneration steps in ketone complexes of borane-oxazaborolidine adducts functioning as chiral catalysts were investigated by using ab initio MO methods. The hydride transfer was found to be highly exothermic. Formation of a novel 1,3-oxazadiboretane structure (e.g., I) was found to precede the regeneration of the catalyst. The regeneration occurring via a cleavage of the 1,3-oxazadiboretane ring was found to require about 101 of the energy released in the hydride transfer. Reactions being potentially involved in the deactivation of oxazaborolidine catalysts were

12/09/2007

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L3 ANSWER 30 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1991:655308 CAPLUS DOCUMENT NUMBER: 115:255308 TITLE: Quentum charia-1
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115:255308 Quantum chemical modeling of chiral catalysis. Part 3. On the role of a Lewis basic solvent in the mechanism of catalytic

enantioselective

reduction of carbonyl compounds by chiral oxazaborolidines

AUTHOR(S): CORPORATE SOURCE:

Oxazaborolidines Nevalainen, Vesa Dep. Chem., Univ. Helsinki, Helsinki, SF-00100, Finland

Tetrahedron: Asymmetry (1991), 2(8), 827-42 CODEN: TASYE3; ISSN: 0957-4166 SOURCE:

DOCUMENT TYPE: Journal

LANGUAGE:

MENT TYPE: Journal UAGE: English English Structures and energies of formation of complexes of Lewis basic solvents with borane-oxazaborolidine adducts functioning as chiral catalysts were investigated by using ab initio MO methods (6-316-'/6-316-'). Formation of complexes of water with a borane adduct of 1,3,2-oxazaborolidine and simpler analogs of it was examined as a model system. Coordination of water to the borane adduct

1,3,2-oxazaborolidine stabilized the adduct by about 50-60% of that of a free borane. Substitution of water bound to the borane adduct of the catalyst by formaldehyde required about 4-5 times more energy than coordination of formaldehyde to the corresponding solvent-free borane

L3 ANSWER 31 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1991:583405 CAPLUS
DOCUMENT NUMBER: 115:183405
TITLE: Quantum chemical modeling of ct Quantum chemical modeling of chiral catalysis. Part 2. On the origin of enantioselection in the coordination of carbonyl compounds to borane adducts of chiral oxazaborolidines
Nevalainen, Vesa
Dep. Chem., Univ. Helsinki, Helsinki, SF-00100,
Finland
Tetrahedron: Asymmetry (1991), 2(6), 429-35
CODEN: TASYE3; ISSN: 0957-4166 AUTHOR(S): CORPORATE SOURCE: SOURCE: CODEN: TASYE3; ISSN: 0957-4166

DOCUMENT TYPE: Journal

AB Energies of formation and structural parameters of HCHO and MeCHO complexes of a model system of borane adducts of oxazaborolidine type of chiral reduction catalysts (CBS reduction) were calculated by using ab initio MO methods

(6-31G*//6-31G*). The energetic preference was determined for the formation of complexes in which the Lewis acidic boron of the borane adduct of an oxazaborolidine would coordinate either syn or anti to the Me group of acetaldehyde. The formation of anticomplex was favored by 15.2 kJ mol-1 which corresponds to a relative anti:syn abundance ratio of 461:1 461:1

and an enantiomeric excess of 99.8%.

L3 ANSWER 32 OF 32 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1991:429427 CAPLUS DOCUMENT NUMBER: 115:29427 TITLE: Quentum chemical modeling of ch

AUTHOR(S): CORPORATE SOURCE:

SOURCE:

DOCUMENT TYPE: LANGUAGE: AB Energies of

SSION NUMBER: 1991:429427 CAPLUS
HENT NUMBER: 115:29427
E: Quantum chemical modeling of chiral
catalysis. On the mechanism of catalytic
enantioselective reduction of carbonyl compounds by
chiral oxazaborolidines

OR(S): Nevalainen, Vesa
DEP. Chem., Univ. Helsinki, Helsinki, SF-00100,
Finland
CE: Tetrahedron: Asymmetry (1991), 2(1), 63-74
CODEN: TASYE3; ISSN: 0957-4166
MENT TYPE: Journal
LINGE: English
Energies of formation and structural parameters of two model systems of
oxazaborolidine type of chiral reduction catalysts (CBS
reduction), their borane adducts, and formaldehyde complexes of the
borane adducts were calculated by using ab initio MO methods. Energies
he

he formation of formaldehyde complexes in which the borane and carbonyl were cis about the B-N bond of the oxazaborolidine ring were found to be slightly pos. The corresponding trans coordination was found to be repulsive. A new class of potential chiral catalysts which also contain the substructure O-B-N was found.

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(WO 2005035540/AP, PRN)

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12/09/2007 10574,871>

TRANSFER L5 1- PN : 9 TERMS

L7 9 L6

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=> S L7 AND US/PC

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